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- Wettable, flexible, oxygen permeable contact lens containing block copolymer polysiloxane-polyoxyalkylene backbone units.
- © Contact lenses which are optically clear, wettable, flexible, and of high oxygen permeability in the aqueous ocular environment of use, of a block copolymer containing polysiloxane and polyoxyalkylene oxide units are disclosed, as well as the preparation thereof and methods of treating vision defects therewith.

Wettable, Flexible, Oxygen Permeable Contact Lens Containing Block Copolymer Polysiloxane-Polyoxyaikylene Backbone Units

This invention relates to ophthalmic devices, such as contact lenses and intraocular implants, and particularly contact tenses of a block copolymer containing polysiloxane and polyoxyalkylene oxide units possessing an advantageous blend of desirable properties including a) high oxygen permeability, b) good wettability, c) flexibility and d) optical clarity, in the ocular environment of use. The invention further relates to the polymers from which the ophthalmic devices are fabricated and the special macromers used to prepare the polymers.

The use of siloxane containing materials of various types in the fabrication of ophthalmic devices is well known.

Thus, in U.S. 3,996,187; 3,996,189; 3,341,490 and 3,228,741 there are described contact lenses fabricated from poly (organosiloxanes) containing fillers. While such lenses are generally soft and of high oxygen permeability, the use of fillers such as silica, is indicated in order to increase the otherwise generally poor tear strength and tensile strength. Also, such silicone rubber lenses are characteristically both hydrophobic and lipophilic.

U.S. 3,808.178 discloses hard contact lenses fabricated from copolymers of a polysiloxanylalkyl acrylate or methacrylate ester and an alkyl acrylate or methacrylate ester. The monomers and polymers of the patent are hydrophobic and are incompatible with hydrophilic materials such as hydroxyethyl methacrylate. While the disclosed tenses have increased oxygen permeability, they are rigid.

U.S. 4.136,250 discloses hydrogels of a copolymer of about 20 to about 90 % of a hydrophilic (or mixture of hydrophilic and hydrophobic) monomer and about 10 to about 80 % of a polyolefinic siloxane macromer which can be used to fabricate a contact lens. Typically, such materials are recited to possess a degree of swelling of about 10 to about 12 %. It has been found, however, that the presence of substantial amounts of water therein limits the oxygen permeability of such materials.

U.S. 4,153,641 relates, in relevant part, to contact lenses fabricated from a polymer of a polyor-ganosiloxane terminated with vinylic groups, or copolymers thereof with other monomers. The exemplified products therein are hydrophobic in nature.

U.S. 4,486,577 relates to copolymers of about 8 to 70 % of a polysiloxane macromer containing at least two vinyl groups and 30-92 % of a monomer which is at least predominantly water insoluble to make polymers useful, for example, as contact lenses.

Generally, such prior art contact lens compositions are either insufficiently hydrophilic in terms of surface wettability to be acceptable to the contact lens profession, even though they may possess high oxygen permeability, or such contact lenses are of acceptable wettability but the hydrophilicity is coupled with water swellability, which tends to limit optimum oxygen permeability.

It is an object of the present invention to overcome these and other disadvantages of the art by providing ophthalmic devices, such as contact lenses and corneal implants, possessing a high degree of surface wettability and possessing a high degree of oxygen permeability notwithstanding their swellability by employing a crosslinked block polymer containing polysiloxane and polyalkylene oxide units.

A further object of the invention is to provide a method of correcting visual defects in the form of refractive errors by fitting to the patient's eye in need of the same a corrective contact tens of such polymer. These and other objects of the invention are apparent from the following detailed description of the invention.

Another object of the invention is to provide novel macromers for accomplishing the above objects.

The present invention is directed to macromers, polymers derived therefrom, ophthalmic devices obtained from such polymers as well as methods of manufacturing each of the foregoing. Each of the above invention materials comprises segments "Seg" of the formula

45 (R·a-Sil-[(L)<sub>h</sub>-R<sub>1c</sub>-Sil]<sub>n</sub>-(L)<sub>h</sub>-R<sub>1s</sub>-A-[L-R<sub>1s</sub>-A]<sub>m</sub>) (I) wherein Sil is a divalent poly(disubstituted siloxane). A is a divalent polyoxyalkylene oxide (which may be substituted). R<sub>1s</sub> is oxygen or an amino nitrogen, R<sub>1a</sub> is a bond, an amino nitrogen, or, when the adjacent atom of Sil is other than silicon, it may also be oxygen, L is a divalent linking group, n is an integer of 0 to 3, and m is an integer of 0 to 10.

The present invention relates to novel macromers, polymers fabricated therefrom, ophthalmic devices prepared from the polymers (especially contact lenses and intraocular lenses), as well as methods of manufacture of each. The ophthalmic devices ultimately fabricated are optically clear, hydrolytically stable, biologically inert, wettable, flexible, of good mechanical strength, oxygen permeable, and when fully swollen in their environment of use have a water content of at least 10 %.

The macromer is a reactive monomer having a segment "Seg" of the formula (R14-Sil-[(L)h-R14-Sil]h-(L)h-R15-A-[L-R15-A]m) more preferably a segment "Grp" of the formula  $\{R_{15}-A-[L-R_{15}-A]_{\overline{m}}-(L)_h\}_{\overline{a}}-(Seg-(L)_h)_{\rho}-Seg-((L)_h-R_{14}-Sil-[(L)_h-R_{14}-Sil]_n\}_{\overline{a}}$ 

and most preferably the monomer is of the formula Cap-Grp-Cap

wherein "Cap", "Grp" and "Sil" are singular variables (i.e. the "a" within "Cap" is not a separate variable); a is zero or 1;

e is zero or 1;

10 a plus e is 1:

h is 1, but may also be zero when the atom of Sil, to which it would otherwise be attached, is a carbon atom:

n is zero to 3;

m is zero to 10, preferably zero to 3;

15 p is zero to 7:

each R14 is a bond, -N(R1)-, or -O- except that R14 cannot be -O- if it is attached directly to a silicon atom; Ris is -O- or -N(R·)-, with Ri being hydrogen, Ci-calkyl, or phenyl; each Sil is independently of the formula

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in which  $R_2$  and  $R_5$  are each independently branched  $C_{3-7}$  alkylene, carbonyl,  $C_{1-6}$  alkylene carbonyl, C.-alkylene-(oxy-C:-alkylene)g- [g being an integer of 1 to 10], or C:-alkylene-O-CH2-(hydroxy C:--«alkylene)-CH2-;

each b and f is independently 0 or 1;

each R3 and R4 is independently selected from C1-18alkyl and aryl of up to 12 carbon atoms; and each y is independently 1-200;

each L is independently selected from -L: R<sub>6</sub>L<sub>2</sub>- wherein L: and L<sub>2</sub> are each independently

-  $\overset{I}{C}$  NH- in which the oxygen or amino nitrogen respectively is bound to  $R_6$ , or independently  $L_1$  and  $L_2$ 

O  $\Gamma$  = C - or a bond, and each  $R_6$  is independently selected from

i) a divalent aliphatic group of up to 25 carbon atoms;

ii) a group of the formula

-(C<sub>1</sub>-20aliphatic)<sub>j</sub>-Q<sub>k</sub>-{(5-7 membered cycloaliphatic or aryl of up to 25 carbon atoms)-Q<sub>(k1)</sub>-(C<sub>1-20</sub>aliphatic)-בפונים

each of j, k, k1 and j1 being independently zero or one, but if j is 0, then k is 0 and if j1 is 0, then k1 is 0; wherein each aliphatic group in i) and ii) above may be interrupted at any point by Q, and each aliphatic, cycloaliphatic and aryl group in i) or ii) may be singly or multiply substituted by one or more of halogen, hydroxy, C1-calkyl, carboxy or C1-12- perhaloalkyl;

Q is -O-, -C-O-, -O-C-, -N-, -N-C-, -C-N-, -OC-N-, -N-CO-, -O-C-O-, 
$$R_1$$
 O  $R_1$   $R_1$  O  $R_1$  or -N-C-N-; and D is 1 to 3;

each A is independently of the formula

wherein each r is 0-4:

t is 3-200, in which preferably at least 30 %, more preferably at least 50 %, still more preferably at least 75 %, most preferably 100 %, of the A groups of formula V within any one macromer of formulae I-III are limited to having homopolymeric sub-blocks of not more than 15 repeating units, preferably not more than 10 repeating units, more preferably not more than 7 repeating units, most preferably not more than 4 repeating units: each R<sub>7</sub> is independently selected from groups i) and iii) below and each R<sub>8</sub> is independently selected from groups i)-iv) below:

i) hydrogen, halogen, unsubstituted  $C_{1-\epsilon}$  alkyl, substituted  $C_{2-\epsilon}$  alkenyl, unsubstituted  $C_{2-\epsilon}$  alkenyl, wherein a) the substituents on the alkyl and alkenyl groups in i) are independently selected from fluoro.  $C_{1-\epsilon}$  alkoxy,  $C_{2-\epsilon}$  alkanoyloxy,  $C_{1-\epsilon}$  alkoxy-C(O),  $C_{2-\epsilon}$  alkenyloxy-C(O),  $C_{3-\epsilon}$  alkenyloxy, aryl of up to 10 carbon atoms, aryloxy of up to 10 carbon atoms, aroyl of up to 11 carbon atoms, aroyloxy of up to 10 carbon atoms, aroyloxy of up to 11 carbon atoms, aryl (of up to 10 carbon atoms)-oxycarbonyl,  $C_{3-\epsilon}$  cycloalkyl,  $C_{3-\epsilon}$  cycloalkoxy,  $(C_{3-\epsilon}$  cycloalkoxy,  $(C_{3-\epsilon}$  cycloalkoxy) carbonyl, oxacycloalkyl of up to 7 carbon atoms, oxacycloalkoxy of up to 7 carbon atoms, oxacycloalkoxy (of up to 7 carbon atoms)-carbonyl, and oxacycloalkyl (of up to 7 carbon atoms)-carbonyloxy; and b) each alkyl and alkenyl group in a) being unsubstituted or further substituted by fluoro,  $C_{1-\epsilon}$  alkyl, or  $C_{1-\epsilon}$  alkoxy provided that said last mentioned alkoxy is not bound to a carbon already singly bound to another oxygen atom; and

ii)  $C_{1-16}$  alkoxy,  $C_{2-16}$  alkanoyloxy,  $C_{1-16}$  alkoxy-C(O)-,  $C_{2-16}$  alkenyloxy-C(O)- and  $C_{3-16}$  alkenoyloxy, each of which may be further substituted by fluoro, anyl of up to 10 carbon atoms, or  $C_{1-6}$  alkoxy provided that the last mentioned alkoxy is not bound to a carbon atom which is singly bound to another oxygen:

(iii) aryl of up to 10 carbon atoms, C<sub>3-8</sub>cycloalkyl, and oxacycloalkyl of up to 7 carbon atoms, each of which is unsubstituted or substituted by a substituent selected from those indicated in i)a) and b) above;

iv) aryloxy of up to 10 carbon atoms, cycloalkoxy of up to 8 carbons, cycloalkyl (of up to 8 carbons)-C(O)O-, cycloalkoxy (of up to 8 carbons)-carbonyl, aroyloxy of up to 11 carbons, oxacycloalkoxy of up to 7 carbons, oxacycloalkoxy of up to 7 carbon atoms)-carbonyl, oxacycloalkyl (of up to 7 carbon atoms)-carbonyloxy, aryloxy of up to 11 carbon atoms, and aroyloxy of up to 11 carbon atoms, each of which is unsubstituted or substituted by fluoro,  $C_1 = 6$  alkyl, or  $C_1 = 6$  alkoxy, provided said last mentioned alkoxy is not bound to a carbon atom which is singly bound to another oxygen; and 2 adjacent groups  $R_7$  and  $R_8$ , together with the atoms to which they are attached may define a 5-8 membered cycloalkyl, cycloalkenyl, or oxacycloalkyl ring or a 6-14 membered bicyclo-ring.

Each Cap is independently selected from
i) a vinyl containing group of the formula  $R \cdot _{5}(L_{4})_{z}(R_{5})_{w}(A \cdot )_{x}(L_{3})_{x}[J(R_{3})_{M}(L_{5})_{G}(R_{5})_{x}(L_{5})_{T}]_{-Z} \qquad (VI)$ or ii) a group of the formula  $R \cdot _{7}(L_{4})_{x}(R_{5})_{w}(A \cdot )_{x}(L_{3})_{x}[J(R_{9})_{M}(L_{5})_{G}(R_{5})_{x}(L_{5})_{T}]_{-Z} \qquad (VII)$ at least one Cap per macromer being of formula VI; wherein  $R \cdot _{5} \text{ is of the formula}$ 

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$$R_{10} = C - C - (VIII)$$

in which  $R_{12}$  is hydrogen or  $C_{1-2}$ alkyl and each of  $R_{10}$  and  $R_{11}$  is independently hydrogen,  $C_{1-2}$ alkyl, or  $-COOR_{13}$  with  $R_{13}$  being hydrogen or  $C_{1-4}$ alkyl;

w. x and z are each independently zero or one but if w is zero, one of x and z must be zero:

with the right hand portion attached to Grp provided that the Grp atom to which  $L_2$  is attached is not a carbonyl carbon, an amido nitrogen or a carboxy oxygen and  $L_3$  is -O- or -NR<sub>1</sub>- when such Grp atom is a carbonyl carbon;

when w is one, and selected from the same group as L<sub>3</sub> when w is zero;

$$A_{1} \text{ is } \begin{array}{c|cccc} & R_{7} & R_{8} & R_{7} \\ & & C & (C)_{7} & C & O \\ & & & & \\ & & R_{7} & R_{8} & R_{7} \end{array}$$

L<sub>5</sub> is -C(0)NH- with the carbonyl attached to R<sub>2</sub> or J;

La is -C(O)NH- with the nitrogen attached to Ra or Ra:

5 J is -O- or -NH-;

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G. Y. M. T. V. and Z are independently 0 or 1, but if Y is zero, then G is zero and M is one; and

$$R_{9} \text{ is } \begin{array}{c|c} R_{7} & R_{8} & R_{7} \\ | & | & | \\ \hline R_{9} & | & | & | \\ R_{7} & R_{8} & R_{7} \end{array}$$

and R-7 is selected from

- a) H, NH<sub>2</sub>, OH, -NCO, -NCS, -C(O)-halogen and COOR<sub>12</sub> when the non-R<sub>17</sub> atom to which it is attached is a non-carbonyl saturated carbon or the non-R<sub>17</sub> atom to which it is attached is a carbon which is part of a ring;
- b) H, -C(O)-halogen, -C(O)-R<sub>12</sub>, -C(O)-OR<sub>12</sub> and R<sub>12</sub> when the non-R<sub>17</sub> atom to which it is attached is an oxygen other than a carboxy oxygen;
  - c) H and R<sub>12</sub>, when the non-R<sub>17</sub> atom to which it is attached is a carboxy oxygen;
- d)  $-C(O)-R_{12}$ ,  $-C(O)-OR_{12}$ ,  $-C(O)N(R_1)R_{12}$  and  $R_{12}$  when the non- $R_{17}$  atom to which it is attached is an amino nitrogen, or  $R_{12}$  when the non- $R_{17}$  atom to which it is attached is an amido nitrogen;
- e) OH, R<sub>12</sub>, CH<sub>2</sub>C(O)R<sub>12</sub>, CH<sub>2</sub>C(O)OR<sub>12</sub>, NH<sub>2</sub>, and -NHR<sub>12</sub> when the non-R<sub>17</sub> atom to which it is attached is a carboxy carbon; and

f) H. OH, halogen, -OR12 when the non-R17 atom to which it is attached is silicon.

Similar macromers and e.g. polymers thereof are disclosed in European Patent Applications 330.616, 330.617 and 330.618. However, the residues  $R_2$  and  $R_3$  of Sil (formula IV) of the present invention are different from the corresponding residues in the above mentioned applications.

The macromers of formulae I, II and III or mixtures thereof are polymerized into polymers which may be shaped into ophthalmic devices such as intraocular implants and contact lenses or utilized as active agent releasing materials, especially release rate controlling membranes or matrices in transdermal drug delivery devices or as tablet release matricies in various applications. The polymers and ophthalmic devices of the invention will swell, when in contact with sufficient aqueous medium, until a fully swollen water content of from at least 10 % to about 95 % has been reached.

The ophthalmic devices and polymers of the invention exhibit a receding contact angle of less than 60; the measurement of which is conveniently performed using a modified "Wilhelmy Plate" technique, as described for example in J.D. Androde et al. Surface and Interfacial Aspects of Blomedical Polymers, Vol. 1,

Surface Chemistry and Physics. Plenum Press, 1985, wherein a specimen sample, in the form of a plate of known dimensions, is immersed into the wetting solution, pure water, at a slow controlled rate, e.g. at 2-20 mm per minute.

The invention ophthalmic devices also have a high oxygen permeability, Dk. Dk is measured using a modification of ASTM standard D3985-81 in that a) 21 % oxygen, i.e. air, is used instead of 99-100 % oxygen, b) 0.5 m² surface area is used instead of 100 m², and c) the humidity is 95-100 % relative humidity instead of 0 % relative humidity. Typically, conventional, fully swollen polyhydroxyethylmethacrylate (pHEMA) lenses which are sparingly crosslinked have a Dk of 5-7 (5-7 x 10<sup>-10</sup> · mm mml O²/cm² sec mmHg) or 5-7 x 10<sup>-10</sup> cm/sec. The ophthalmic devices of the instant invention have oxygen permeability Dk values generally greater than 7-10 (10<sup>-10</sup> cm/sec).

Preferred macromers of formulae I, II and III are those wherein n is zero or one, most preferably zero. Another group of preferred macromers of formulae I-III are those where m is 0-4, more preferably zero or one, most preferably zero. A highly preferred group of macromers of formulae I-III are those wherein both m and n are zero or one, most preferably both m and n are zero, to result in Seg., a most highly preferred embodiment of Seg. of the formula

 $\{R_{14}-Sil-(L)_h-R_{15}-A\}$  (la)

and Grp1, a most highly preferred embodiment of Grp, of the formula

 $\{R_{15}-A-(L)_h\}_a-Seg_1-((L)_h-R_{14}-Sil\}_e$  (IIa).

Within Sil above, b and f are each independently preferably one, with a highly preferred embodiment being when both b and f are one;

 $H_2$  and  $H_3$  are each independently preferably branched  $C_{2-7}$  alkylene,  $C_{1-4}$  alkylene-(oxy- $C_{2-4}$  alkylene)<sub>0</sub>-(wherein g is 1-10, preferably 1-4, more preferably 1-2, most preferably 1) or  $C_{2-4}$  alkylene-oxy- $C_{1-2}$  alkylene)- $C_{1-2}$  alkylene)- $C_{1-4}$  alkylene or  $-C_{2-4}$  with g preferably 1 or 2, most preferably 2.

Within Sil of formula IV, y is preferably 5 to 100, more preferably 10 to 50, most preferably 20 to 30. A particularly useful subembodiment has y = 25 to 29, especially when a is 1 and e is zero.

R14 is preferably a bond or -O-, more preferably -O-.

R<sub>1</sub> is preferably methyl or hydrogen, more preferably hydrogen.

Each R<sub>3</sub> and R<sub>4</sub> is independently preferably C<sub>1-7</sub>alkyl or phenyl, more preferably C<sub>1-4</sub>alkyl, still more preferably methyl or ethyl, most preferably methyl. A most preferably embodiment of Sil is when all of R<sub>3</sub> and R<sub>4</sub> are the same.

R<sub>1s</sub> is preferably -O- or -NH-, more preferably -O-.

Each h is independently preferably 1, most preferably each h is one.

p is preferably 0 to 5, still more preferably 0 to 3, most preferably 0 or 1.

As stated above, the linking group L is  $L_1R_6L_2$  wherein  $L_1$  and  $L_2$  are independently -COO- or -C(O)NH- (in which the oxygen or amido nitrogen is bound to  $R_6$ ) or -C(O)- or a bond. Preferably, each  $L_1$  and  $L_2$  is independently -COO-, -C(O)NH-, or -C(O)-, more preferably -COO-, or -C(O)NH- and most preferably -C(O)-NH-.

In a most highly preferred embodiment, all of the L<sub>1</sub> groups are the same and all of the L<sub>2</sub> groups are the same. In a still more highly preferred subembodiment, L<sub>1</sub> and L<sub>2</sub> are the same groups.

Each R<sub>6</sub> is preferably independently selected from

- i) a divalent aliphatic group of up to 12 carbon atoms, preferably up to 7 carbon atoms, more preferably up to 4 carbon atoms;
- ii) -(C<sub>1-20</sub>aliphatic)<sub>I</sub>-Q<sub>k</sub>-((5-7 membered cycloaliphatic or aryl of up to 25 carbon atoms)-Q<sub>k1</sub>-45 (C<sub>1-20</sub>aliphatic)<sub>I</sub><sub>1</sub><sub>D</sub>- each of j, k, k1 and j1 being 0 or 1 but if j is zero, k must be zero and if j1 is zero, k1 must be zero: preferably each of the C<sub>1-20</sub>aliphatic groups in ii) are C<sub>1-12</sub>aliphatic, more preferably C<sub>1-2</sub>aliphatic, still more preferably C<sub>1-4</sub>aliphatic, most preferably C<sub>1-2</sub>aliphatic; preferably the carbons is up to 14 carbons, more preferably up to 10 carbons, still more preferably of 6 carbons, most preferably a phenyl ring; preferably k is zero and preferably k1 is zero, but most preferably k and k1 are each zero; wherein each aliphatic group within i) and ii) above may be interrupted at any point by Q and each aliphatic, cycloaliphatic, and aryl group in i) or ii) is unsubstituted or substituted by one or more of halogen (preferably fluorine or chlorine, most preferably fluorine), hydroxy, C<sub>1-4</sub>alkyl (preferably methyl, 12perhaloalkyl (preferably C<sub>1-7</sub> perhaloalkyl, more preferably C<sub>1-4</sub>perhaloalkyl, still more preferably perhalomethyl, perhaloethyl, or perhalopropyl, most preferably perhalomethyl, with the halogen atom being preferably the same and preferably fluorine or chlorine, most preferably fluorine. D is preferably 1 or 2, more preferably 1 or 2.

Most preferably, the R<sub>6</sub> aliphatic groups are not interrupted by Q, and preferably are unsubstituted. Also most preferably, the R<sub>6</sub> cycloaliphatic groups are substituted by 1-4 substituents, preferably 1-3 substituents, which substituents are preferably selected from halogen (preferably fluorine), hydroxy, and C·-calkyl (preferably methyl, ethyl, propyl or isopropyl, most preferably methyl). In a most highly preferred embodiment each of the substituents is the same. Preferably, one of j and j1 is zero when the R<sub>6</sub> group contains a cycloaliphatic. The most highly preferred cycloaliphatic containing R<sub>6</sub> is (5-yl-1,3,3-trimethyl-cyclohexyl)-methyl (the residue of isophorone diisocyanate absent the two isocyanate groups).

Preferably, the R<sub>s</sub> aryl groups are unsubstituted or substituted by 1-4 substituents, preferably unsubstituted or substituted by 1 or 2 substituents, more preferably unsubstituted or substituted by 1 substituent which substituents are preferably selected from halogen (preferably fluorine), hydroxy, and C<sub>1-4</sub>alkyl, more preferably C<sub>1-4</sub>alkyl (preferably methyl, ethyl, propyl, or isopropyl, more preferably methyl). When more than one substituent is present per aryl, the substituents on any one aryl is preferably the same. A most highly preferred aryl embodiment of R<sub>s</sub> is toluenediyl.

A preferred subclass of R<sub>6</sub> includes: ethane-1,2-diyl; propane-1,2-diyl; propane-1,3-diyl; tetramethylene-diyl; hexane-1,6-diyl; cyclohexane-1,2-diyl; cyclohexane-1,3-diyl; 1,4-phenylene; bis[4-yl-cyclohexyl]-methane; bis(4-yl-cyclohexenyl)methane; bis(4-yl-phenyl)methane; toluene-2,4-diyl; toluene-2,6-diyl; m- and p-tetramethylene-diyl; toluene-1,2-diyl; toluene-1,4-diyl; 3,3'-dichloro-4,4'-diyl-biphenyl; naphthalene-1,5-diyl; diphenyl; tetrahydrotoluene-diyl; (5-yl-1,3,3-trimethylcyclohexyl)methyl; trimethylhexane-1,6-diyl; (especially 2,2,4-trimethylhexane-1,6-diyl and 2,4,4-trimethylhexane-1,6-diyl); diethylfumarate-2,2'-diyl; 1-carboxypentane-1,5-diyl; naphthalene-1,2-, 1,3-, 1,6-, 1,7-, 1,8-, 2,7, and 2,3-diyl; 1-methylnaphthalene-2,4-diyl; methylcyclohexane-1,4-diyl; 6,(7)-methylnaphthalene-1,3-diyl; biphenyl-4,4'-diyl; -3,3'-dimethoxybisphenyl-4,4'-diyl; 2,2'-dimethylbiphenyl-4,4'- and 3,3'-diyl; bis (4-ylphenyl)ethane; and bis(4-ylphenyl)ether.

The polyalkylene oxide groups are of formula V and are homo or copolymeric blocks having a total of 3-200 alkylene oxide units and an additional terminating alkylene oxide or amine. However, no homopolymeric subblock is in excess of 100 repeating units. Preferably, no individual A has a subblock in excess of 15 repeating units. The portion of the alkylene oxide alkylene chain in the macromer backbone is a straight chain of 2 (r is zero) to 6 (r is 4) carbon atoms which may be unsubstituted or substituted as set forth previously.

Preferably the number of alkylene oxide units (i.e. t) is 3-150, more preferably 4-100, still more preferably 5-75, still even more preferably 6-70. Within this group are two highly preferred subembodiments, a) those with t of 25-70, preferably 50-68, most preferably 60-66 and b) those with t of 8-50, preferably at least 9, more preferably at least 10, still more preferably 13-35, even more preferably 15-30, most preferably 20-25, especially about 22.

Also highly preferable are two embodiments wherein at least 30 % of the A groups within the macromers of formulae I-III consist of 32 % and 35 %, respectively, of polyethylene glycol and 68 % and 65 %, respectively of polypropylene glycol, the polyethylene glycol blocks being of no more than 3 repeating units each and the polypropylene glycol blocks being of no more than 7 repeating units each.

While it is highly preferable that each A be of the same structure throughout the macromer, there is no such absolute requirement. Preferably at elast 20 %, more preferably 25 %, still more preferably 33 %, yet more preferably 50 %, even more preferably 75 % and most preferably at least 90 % of the A groups within any one macromer are the same.

Within each A group, there are two highly preferred subembodiments:

- a) homopolymeric A units,
- b) copolymeric A units of the 'EXE' type with the 'X' portion having approximately twice the number of repeating units as either 'E' (i.e. a poloxamer or reverse poloxamer like grouping).

While these two arrangements are highly preferred, other arrangements of alkylene oxide copolymeric type are also suitable.

Substituents on the alkylene oxide backbone portion (i.e. R<sub>7</sub> and R<sub>8</sub>) are preferably selected from those set forth above. Preferably, each alkyl group within R<sub>7</sub> or R<sub>8</sub> whether alone or as a part of a larger group, has up to 4 carbon atoms, more preferably methyl, ethyl, butyl, or t-butyl (provided that adjacent groups on a ring cannot each be t-butyl), most preferably methyl. Preferably, each alkenyl within R<sub>7</sub> or R<sub>8</sub>, whether alone or as part of a larger group has 2-4 carbons, and is more preferably C<sub>2-3</sub>alkenyl. An R<sub>7</sub> or R<sub>8</sub> aryl is preferably phenyl or naphthyl, most preferably phenyl. Cycloalkyl groups within R<sub>7</sub> and R<sub>8</sub> preferably have 5-6 ring members, most preferably 6 ring members. Oxacycloalkyl groups within R<sub>7</sub> or R<sub>8</sub> are preferably of 5-6 ring members, more preferably 6 members, each having only a single heteroatom in the ring which

While the entire A unit may have a substantial fluorine substituent content, it is preferably at least 25 %

halogen free, more preferably at least 30 % halogen free, still more preferably at least 40 % halogen free, yet more preferably at least 60 % halogen free, and most preferably substantially halogen free.

Preferably R<sub>7</sub> is hydrogen, C<sub>1-7</sub>alkyl more preferably C<sub>1-4</sub>alkyl, more preferably methyl; substituted C<sub>2-4</sub>alkyl, most preferably substituted methyl; phenyl, substituted phenyl, benzyl, substituted benzyl, cyclohexyl, or C<sub>4-5</sub>oxacycloalkyl, preferably C<sub>5</sub>-oxacycloalkyl; each of the "substituents" preferably being C<sub>2-5</sub>alkoxy (more preferably C<sub>2-6</sub>alkoxy), fluorine, or when substituted on a group other than alkyl, C<sub>2-5</sub>alkyl; the most preferable substituents being methoxy, fluorine, and when substituted on other than alkyl, methyl. Substituents on aryls within R<sub>7</sub> are preferably in the p-position. R<sub>8</sub> is preferably selected from the same group as R<sub>7</sub> and C<sub>1-4</sub>alkoxy, phenoxy, and C<sub>5</sub>cycloalkoxy, more preferably the same group as R<sub>7</sub>.

At least one Cap per macromer is independently of formulae VI or VII, but preferably the macromers have both Cap groups of formula VI. Most preferably, within each macromer the Cap groups are the same.

Preferably, the Cap groups are of formula VI in which R<sub>16</sub> is preferably a vinyl group of formula VIII in which R<sub>10</sub> is hydrogen, R<sub>10</sub> is hydrogen, C<sub>1-4</sub>alkyl (preferably methyl), or -COOR<sub>13</sub>, with R<sub>10</sub> being hydrogen or methyl. More preferably the formula VI Cap groups have an R<sub>16</sub> which is a vinyl of formula VIII in which R<sub>10</sub> and R<sub>10</sub> are each hydrogen and R<sub>12</sub> is hydrogen or methyl.

Preferred Cap groups, whether of formula VI or VII are those wherein

a) Z, w. x. z and V are all zero;

b) Z, w, V, and x are all zero, z is one, and L<sub>z</sub> is -C(O)-, -C(O)NH-, or -C(O)O-, most preferably -C(O)-O, or -C(O)- with the carbonyl in each case attached to  $R_{15}$  or  $R_{17}$ :

c) Z, z and V are all zero, w and x are each one,  $R_6$  is of formula IX wherein j, k, j1 and k1 are each zero, and D is one, and the aryl thereof is phenyl; and  $L_3$  is -NHC(O)- with the nitrogen thereof attached to  $R_6$ :

d) Z, w and x are zero, and V is one, A· is  $-CH_2CH_2O$ - with the carbon thereof attached to L<sub>4</sub>, L<sub>4</sub> is -C(O)O- with the carbonyl thereof attached to R<sub>16</sub> or R<sub>17</sub>;

e) Z. V and x are zero, z and w are one, L is -C(0)0- with the carbonyl attached to  $R_{15}$  or  $R_{17}$ , and  $R_{6}$  being alkylene or hydroxy substituted alkylene of from 2-15 carbon atoms; and

f) those groups of a) to e) above except that Z is one and wherein

i) M and T are one, G and Y are zero.  $R_3$  is  $C_{2-15}$ alkylene or  $C_{2-15}$ hydroxy substituted alkylene and  $L_5$  is -NHC(0)- with the nitrogen attached to  $R_6$ :

ii) M, G, Y and T are all one,  $R_3$  is  $C_{2-15}$ alkylene or  $C_{2-15}$ hydroxy substituted alkylene,  $L_5$  is -NHC(O)O- with the oxygen bound to  $R_3$ ,  $R_6$  is as defined above, preferably a divalent aliphatic or cycloaliphatic or arylene group and  $L_6$  is -NHC(O)- with the nitrogen attached to  $R_6$ ;

iii) M is zero, G, Y and T are all one and  $L_5R_5L_6$  is -C(O)NH-R6-NHC(O)-.

Even more highly preferred macromers of formulae I-III are those within formula Ia and IIa having each Cap independently selected from formulae VI and VII in which Z is zero to result in Cap of the formulae  $R_{-5}(L_2)_z(R_5)_w(A_1)_w(L_3)_x$  (VIa) and

 $R \cdot r(L_4)_z(R_6)_w(A_1)_v(L_3)_{\overline{x}}$  (VIIa)

and macromers of the formula

Sil is

Cap,-Grp,-Cap. (IIIa)

Very highly preferred macromers of formula Illa have R<sub>16</sub> in formula VIa selected from those of formula VIII wherein each of R<sub>10</sub>-R<sub>12</sub> is independently hydrogen or methyl. Also preferred within this very highly preferred group are those wherein V is zero.

Highly preferred vinyl containing Cap groups terminating the macromer are acryl, methacryl, and styryl groups bound directly to or through an ethyloxy or oxyethyloxy to the balance of Cap or to the non-Cap portion of the macromer.

Specifically preferred embodiments of the present invention within formulae IIa and III include those wherein

$$\begin{array}{cccc} \text{CH}_3 & \text{CH}_3 & \text{CH}_3 \\ & & &$$

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e is zero;

Ris and Ris are each -O-;

L is

O CH<sub>3</sub> CH<sub>3</sub>
-CNHCH<sub>2</sub> CH<sub>3</sub>
HNC-

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h is one;
A is (CH<sub>2</sub>CH<sub>2</sub>O)<sub>1</sub>; and
p is zero;
m is zero;

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Further preferred is a macromer of the formula Cap-Sil-L-O-A-L-Sil-Cap

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wherein Sil is 
$$-CH_2CH_2 - (Si-O)_y - Si-CH_2CH_2$$
.

 $CH_3 CH_3$ 
 $CH_3 CH_3$ 

wherein y is 1 to 200. (especially y of about 4, 7, and about 16);

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with the phenyl ring thereof bound to Sil, m is zero to 3; A is -( $CH_2CH_2O$ )<sub>t</sub>- ,

· ^

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wherein t is 3 to 200, or a poloxamer or reverse poloxamer having t polyoxyalkylene units per A group (especially t of about 6-8, about 45-55, and about 90-95); and

In the foregoing, all alkyl groups whether mentioned alone or as part of another group are preferably C·-C<sub>4</sub> alkyl, such as methyl, ethyl, propyl and butyl, especially t-butyl, with the exception that adjacent groups on aryl rings cannot each be t-butyl. These alkyl groups may be straight chain or branched chain. When the alkyl is a substituent on a phenyl ring, it is preferably attached at the para position. Preferably alkenyl groups, whether alone or as part of another group, are C<sub>2</sub>-C<sub>4</sub> alkenyl, such as ethenyl, propenyl and butenyl. Preferred aryl groups (whether alone or as part of another group) are phenyl and naphthyl, more preferably phenyl. Preferably the aryl groups are still further substituted by C<sub>1</sub>-C<sub>4</sub> alkyl, more preferably t-butyl, most preferably in the para position. Halogen may be chloro, bromo, iodo or fluoro.

Color may be added to the polymerized material or to the macromers via typical dying techniques. One such manner is by insuring hydroxy groups are introduced onto the macromer backbone or substituent thereof.

Typical color groups useful in these embodiments include, but are not limited to, the hydroxy-reactive dyes known in the art under the tradename Remazol, manufactured by American Hoechst. Examples of the Remazol dyes which are especially useful are:

Dye	Color Index Code
Remazol Brill Blue RW Remazol Yellow GR Remazol Black B Remazol Golden Orange 3GA Remazol Turquoise P	Reactive Blue 19 Reactive Yellow 15 Reactive Black 5 Reactive Orange 78 Reactive Blue 21

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all of which have at least one group of the formula -SO<sub>2</sub>-CH<sub>2</sub>CH<sub>2</sub>O-SO<sub>3</sub> 
which reacts with the polymer or monomer hydroxy group to yield a dye-SO<sub>2</sub>-CH<sub>2</sub>-CH<sub>2</sub>-O-polymer or

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group, preferably the former. In such a manner, both excess free hydroxy groups are disposed of and colored contact lenses can be realized simultaneously.

If the polymerized macromers set forth above have inherently too low a water content, i.e. below 10 % water content, the polymer is not within this invention. However, the macromers can be copolymerized with a hydrophilic modifier so as to increase the water content to be within the instant invention. Whether the hydrophilic modifier is needed or not, and the amount which is needed, is dependent upon the desired polymer water content. Preferably, when the hydrophilic modifier is present, in accordance with the above, it is present in an amount of up to about 50 % by weight preferably about 1 % to about 30 %, more preferably about 5 % to about 25 %, still more preferably about 10 % to about 15 % of the resultant polymer.

In addition to the hydrophilic modifier, other comonomers which may be present in the polymer are monomers coreactive with the monomer of formula I. Such additional monomers may be present in a minor amount of up to about 20 % by weight of the resultant polymer. When such comonomers are excessively hydrophobic, additional hydrophilic modifier may be incorporated so as to achieve the appropriate water content.

The hydrophilic modifier is a monomer which is coreactive with a monomer of formula I and Is typically selected from

aa) polyethylene glycols of the formula

ab) pyrrolidones of the formula

wherein R<sub>19</sub> is hydrogen or C<sub>1</sub>-C<sub>7</sub> alkyl, preferably C<sub>1</sub>-C<sub>4</sub> alkyl, n1 is an integer of from 1 - 25, or

ac) a compound of the formula

 $(R_{20}\text{-div})_{nz}Hy$  (C)

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wherein R<sub>18</sub> is a polymerizable or reactive molety selected from i) ethylenically unsaturated radicals, preferably vinyl, 1-methylvinyl, 2-methylvinyl, or allyl, ii) anhydrides, iii) amines, iv) acids, v) esters, vi) amides, vii) ethers, viii) acid halides, R<sub>20</sub> is selected from the meanings given for R<sub>18</sub> and ix) epoxy groups, especially glycidyl, x) isocyanates and isothiocyanates and xi) hydroxy; div is selected from

ba) a divalent aliphatic group of up to 25 carbon atoms, preferably alkyl, alkenyl or alkynyl, which may also be interrupted, or terminated, or interrupted and terminated by oxy, carbonyloxy, amino, aminocarbonyl, oxycarbonyl, ureido, oxycarbonylamino, or carbonylamino;

bb) a divalent 5 - 7 membered cycloaliphatic or 5 - 7 membered cycloaliphatic-C<sub>1</sub>-C<sub>20</sub>-aliphatic group which may also be interrupted, terminated, or interrupted and terminated as in group ba) above except that said interruptions cannot occur within said cycloaliphatic portions;

bc) a divalent arylene group having 6-25 carbon atoms which is unsubstituted or substituted by at least one substituent selected from halogen, C<sub>1</sub>-C<sub>4</sub> alkyl, and C<sub>1</sub>-C<sub>12</sub> perhaloalkyl;

bd) a divalent aralkyl or alkaryl or -alkylene-aryl-alkylene- having 7 - 25 carbon atoms which is uninterrupted or interrupted in the alkyl portion, or terminated, or interrupted in the alkyl portion and terminated with an interrupting or terminating group as mentioned in ba) above, and each of said uninterrupted, interrupted, and terminated aralkyl and alkaryl groups is further unsubstituted or substituted by a substituent selected from halogen, C<sub>1</sub>-C<sub>4</sub> alkyl, and C<sub>1</sub>-C<sub>12</sub> perhaloalkyl;

wherein na is an integer of 8-100; and

bea) Ra is hydrogen, Rb is methyl, and nb is zero; or

beb) Ra is methyl. Rb is hydrogen, and nb is zero; or

bec) Ra and Rb are hydrogen and nb is one; nz is an integer of 1 up to the valence of Hy and where nz is less than the valence of Hy, the remaining valences of Hy are taken up with hydrogens; and Hy is a hydrophilic group selected from

ca) morpholino; cyclic amide radicals of 5 - 7 ring members; saturated and unsaturated cyclic N,N-diamide radicals of 5 - 6 ring members; groups of the formula

having 6 - 7 ring members wherein nc and nd are selected from 0 - 2; and cyclic amines of 5 - 6 ring members; each unsubstituted or substituted by hydroxy-C<sub>1</sub>-C<sub>5</sub> alkyl, carboxy, or lower alkyl;

cb) tetrahydrofurfuryl;

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- cc) mono-, di-, and polysaccharide radicals, whether straight chain or cyclic, their corresponding sugar alcohol radicals, pentaerythritol radicals, and polyvinyl alcohol radicals; and
- cd) polyhydroxy C<sub>2</sub>-C<sub>7</sub> alkyl radicals; and said reactive group R<sub>18</sub> is capable of reacting with one or more sites in said formula III.

Preferably the hydrophilic modifier has R<sub>20</sub> selected from

ia) 
$$H_2C=C$$
-

and the terminus of div to which it is bound is -C(O)R<sub>25</sub>, the carbonyl being bound to R<sub>18</sub>, where R<sub>25</sub> is hydrogen or methyl and R<sub>26</sub> is -O-, or -NR<sub>27</sub>- with R<sub>27</sub> being hydrogen or lower alkyl;

ib) H<sub>2</sub>C = CH- and the terminus of div to which it is bound is

the left hand bond being bound to  $R_{20}$ , wherein  $R_{28}$  is -O-, -NR<sub>27</sub>-, -C(O)-, -C(O)NR<sub>27</sub>- or -C(O)O-; and

ic) 
$$H_2C=C$$
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- and the terminus of div to which it is bound is -C(O)R<sub>30</sub>-, the carbonyl being bound to R<sub>20</sub> wherein R<sub>30</sub> is lower alkylene or -[CH<sub>2</sub>CH(loweralkyl)-O]<sub>1-5</sub>- and R<sub>29</sub> is hydrogen, lower alkyl, or cyano; and has Hy selected from
  - ce) morpholino which is unsubstituted or mono- up to tetra-substituted by lower alkyl, preferably methyl;

wherein ne is one, two, or three, preferably 1, each of which is unsubstituted or substituted, preferably mono- or disubstituted, by lower alkyl, preferably methyl or ethyl;

$$cg) - N \longrightarrow (CH_2)_{nf}$$

where nf is 2 or 3, preferably 2, each of which is unsubstituted or substituted by lower alkyl, preferably methyl or ethyl; ch) a divalent group of the formula

wherein R31 is hydrogen or carboxy;

ci) 
$$-N$$
 , cj)  $-N$   $(CH_2)_{ng}$ 

where ng is one or two, each of which is unsubstituted or substituted by lower alkyl, preferably methyl or ethyl;

cl) a polyol selected from polyvinyl alcohol radicals.

$$O-CH_2-(CH)_{\overline{xa}}-CH_2O-(CH)_{xb}$$

wherein xa is 2 - 10, preferably 3 - 4, and xb is an integer from zero, preferably at least 0.5 times xa up to xa + 1 inclusive, the cyclic saccharide analogs thereof, the di-tetra saccharides thereof, preferably having saccharide-saccharide links between the 1 and 4 carbon positions thereof, preferably 6-linkages, C(CH2O)4-(H)<sub>xc</sub> wherein xc is 0 - 3, and polyhydroxy lower alkylene glycol radicals wherein up to 50 %, preferably up to 25 %, more preferably up to 10 %, of the hydroxy groups are missing their alcoholic hydrogens.

Most preferably the hydrophilic modifier of formula (C) is selected from

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wherein 
$$R_{32}$$
 is -[OCH<sub>2</sub>(CH)<sub>yc</sub>CH<sub>2</sub>O]-(H)<sub>yb</sub>

in which yc is 1-6, preferably 3-4 and yb is 0 up to (yc + 1), preferably (yc-1) to (yc + 1), more preferably yc to (yc + 1), and ya is 1 up to (yc + 2-yb),

$$CH_3$$
 O O acd)  $[H_2C=C---C-C-CH_2CH_2NHC-]_{yd}$   $R_{33}$ 

wherein  $R_{33}$  is  $C(CH_2O)_{4}(H)_{4,yd}$  and yd is 1 - 4, preferably 1 - 2;

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ace) polyvinyl alcohol having at least one and up to 50 % preferably 25 %, more preferably 10 %, inclusive, of the alcoholic hydrogens thereof absent;

Preferably, when Cap is a vinyl containing group or one or more of the aforementioned R groups within formulae I-III contain an ethylenically unsaturated group, then the macromer can be crosslinked in the presence of a vinylic group containing hydrophilic modifier and/or vinylic group containing crosslinking agents. The instant macromer can be crosslinked in the presence or absence of up to less than about 50%, preferably up to about 30%, most preferably up to about 10% by weight of vinylically unsaturated hydrophilic modifiers and/or other vinylic comonomers.

When Cap does not have a vinylic group, but takes part in crosslinking, Cap is of formula VII. Crosslinking is then typically carried out by condensation with a tri- or polyfunctional coreactive monomer. For example, when Cap terminates in OH, then the coreactive monomer functional group can be -COOH, SCN, -NH<sub>2</sub>, -NH(alkyl) etc.; when Cap terminates in -NH<sub>2</sub> or -NH(alkyl), the reactive comonomer functional group can be a conventional acyl leaving group; and when Cap terminates in OCN- or SCN-, then the reactive comonomer functional group can be OH. Similarly, the other coreactive functional groups mentioned in terms of either Cap or the coreactive monomer can be interchanged.

Suitable vinylic monomers and coreactive monomers for condensation are set forth below. However, the list is not exhaustive and those of ordinary skill will appreciate the modifications, additions, and alternatives which may also be employed.

However, the degree of crosslinking in the finished crosslinked polymer should not exceed 20 %, preferably not greater than 10 %, more preferably not greater than 5 %, still more preferably 1-4 %, most preferably 2-3 %.

A highly preferred embodiment within the polymers fabricated from macromers of formula 1, requires that not more than 20-70 %, preferably not more than 50 % of the A groups are polyethylene glycol. In another highly preferred group A is homopolymeric polyethylene oxide or polypropylene oxide.

The number of A units and the chain length and nature of the substituents on the polyoxyethylene segment is determined by the degree of wettability desired in the polymer of the compound of formula I. In general, the polymer should be sufficiently hydrophilic in its surface properties such that the polymer exhibits a contact angle with distilled water at 20°C of less than 60°, preferably less than 40°, more preferably less than 25°, still more preferably less than 15°, most preferably less than 10°.

The greater the number of siloxane units, the greater the number of oxyalkylene units is generally required to reduce the contact angle to within the above limits.

As water is taken up, the oxygen permeability of the polymers tends to be reduced. However, with the water content and swellability increased due to hydrophilic modifier the drop in Dk over the low water content analogs is surprisingly less drastic than would otherwise be expected. Advantageously, the instant polymers absorb at least 10 % by weight water, preferably 15 - 90 % by weight, more preferably 20 - 85 % by weight, still more preferably 25 - 75 % by weight, most preferably 35 - 55 % by weight water.

Highly preferred are those polymers which exhibit a contact angle of less than 25°, more preferably less than 10°.

The reactive vinylic macromers of the invention can characteristically be polymerized to form crosslinked polymers under conventional polymerization conditions.

If desired, the monomer reaction mixture may contain a catalytic amount of a conventional polymerization catalyst, preferably a free radical catalyst. Of particular interest are conventional peroxide and azo catalysts, such as hydrogen peroxide, benzoyl peroxide, tert-butyl peroctoate, benzoyl peroxide or azobis-(isobutyronitrile).

The polymerization can generally be carried out at temperatures between about 20° and about 150°C, for a period between about 1 and about 24 hours. It is understood that the time and temperature in such a reaction are inversely related. Thus, temperatures employed in the upper end of the temperature range will generally provide reaction times near the lower end of the time range. Preferably, the polymerization is conducted in the presence of actinic radiation, such as UV light.

Depending upon the nature of the polymer mixture, it may be desirable for the polymers obtained from such polymerizations to be post cured, e.g. at a somewhat elevated temperature such as between about 60 °C and about 150 °C.

For the preparation of contact lenses, the polymer mixture may be cast directly in the shape of the lens, or the polymerization may be carried out in a mold having a shape convenient for further processing, such as in the shape of small cylinders or "buttons", which can then be machined.

Minor amounts i.e. less than 50 %, preferably up to 30 %, and most preferably up to no more than about 10 % by weight, of conventional copolymerizable vinyl monomers other than hydrophilic modifiers, can be employed as extenders or the like, in the preparation of the instant polymer, as copolymer constituents. However, if the total hydrophilic modifier content is within the foregoing limits, the hydrophilic modifiers can also be used as such extenders. Suitable vinyl monomers include: acrylates and methacrylates of the general formula

# H<sub>2</sub>C=C-COOR<sub>35</sub>

where R<sub>34</sub> is hydrogen or methyl and R<sub>35</sub> is a straight chain or branched allphatic, cycloaliphatic or aromatic group having up to 20 carbon atoms which is unsubstituted or substituted by one or more alkoxy, alkanoyloxy or alkyl of up to 12 carbon atoms, or by halo, especially chloro or preferably fluoro, or C<sub>3</sub>-C<sub>5</sub> polyalkyleneoxy of 2 to about 100 units; acrylamides and methacrylamides of the general formula

R<sub>34</sub> | H<sub>2</sub>C=C-CONHR<sub>35</sub>

where R<sub>34</sub> and R<sub>35</sub> are as defined above; vinyl ethers of the formula H<sub>2</sub>C = CH-O-R<sub>35</sub>
where R<sub>35</sub> is as defined above; vinyl esters of the formula H<sub>2</sub>C = CH-OOC-R<sub>35</sub>
where R<sub>35</sub> is as defined above;
maleates and furnarates of the formula R<sub>35</sub>OOC-HC = CH-COOR<sub>35</sub>
where R<sub>35</sub> is as defined above;
and vinylic substituted hydrocarbons of the formula
R<sub>34</sub>CH = CHR<sub>35</sub>

where R34 and R35 are as defined above.

Useful monomers include, for example:
methyl-, ethyl-, propyl-, isopropyl-, butyl-, ethoxyethyl-, methoxyethyl-, ethoxypropyl-, phenyl-, benzyl-, cyclohexyl-, hexafluoroisopropyl-, or n-octyl-acrylates and -methacrylates as well as the corresponding acrylamides and methacrylamides; dimethylfumarate, dimethylmaleate, diethylfumarate, methyl vinyl ether, ethoxyethyl vinyl ether, vinyl acetate, vinyl propionate, vinyl benzoate, acrylonitrile, styrene, alpha-methyl styrene, 1-hexene, vinyl chloride, vinyl methyl ketone, vinyl stearate, 2-hexene and 2-ethylhexyl methacrylate.

In one preferred aspect, the instant polymers are free from copolymer units of such conventional monomers which are not hydrophilic modifiers.

The vinylic monomers of formulae I-III can be prepared by methods known, per se.

For example, the siloxane/polyalkylene oxide containing divinylic monomers of formulae I-III may be prepared by reacting a siloxane diol of the formula HO-Sil-H, wherein Sil is a group of formula IV, as defined hereinbefore, with a sufficient amount of a difunctional reactive group containing compound at least one of which reactive groups is coreactive with OH, such as isocyanate; activated carboxy, such as an anhydride, an acid halide or a carboxy ester, or is a leaving group, such as a halide, sulfato or the like, to forming preferred macromers include toluene diisocyanate, isophorone diisocyanate, and polyalkylene oxide mono- or di-carboxy by isocyanate.

The resulting endcapped siloxane derivative can then be reacted with a polyoxyalkylene diol of the formula HO-(A)-H, wherein -A- is a group of formula V above to form the corresponding polyoxyalkylene - siloxane - polyoxyalkylene diol. This diol can be then reacted with a reactive end Cap group containing monomer wherein the reactive group is an isocyanate; activated carboxy, such as an anhydride, an acid halide or carboxy ester, or is a leaving group such as halo, sulfato or the like to form the corresponding difurctions reactive group containing compound having the group -L- to form the corresponding di-functional reactive endcapped derivative which is then reacted with a siloxane of the formula HO-Sil-H. One may continue building up alternative polyoxyalkylene/siloxane unit containing diols in this manner, as desired. Then this diol may be endcapped with a reactive group containing vinylic monomer having a terminal 'Cap' moiety or the diol reacted with a sufficient amount of difunctional reactive group containing compound to endcap the diol with a reactive group, such as e.g. an vinylic amine or alcohol, or other copolymerizable monomer having a crosslinkable group to obtain the corresponding product of formula III.

Of course, instead of starting with a siloxane of the formula HO-Sil-H and building up the alternating sequence therefrom, one may instead begin with a polyoxyalkylene diol of the formula HO-(A)-H and, after endcapping the same with difunctional reactive groups, condense the same with the siloxane until the desired number alternating siloxane/poly alkylene oxide groups is attained and terminate the diol with Cap groups as described above.

Alternative synthesis of the direactive siloxane can be accomplished with a siloxane dihydride of the formula

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and reacting it (as known in the art) with a vinyl containing compound containing another reactive group, typically a hydroxy reactive group, such as styrene isocyanate, to yield the dicapped siloxane, such as

$$\begin{array}{c|c}
R_3 & R_3 \\
\hline
(SiO) - Si \\
R_4 & R_4
\end{array}$$

Such reactive group capped siloxane may then be used in the same manner as the reactive endcapped HO-Sil-H indicated above.

Synthesis of the macromers may be accomplished neat or in solvents as indicated in the examples attached hereto in accordance with known synthetic techniques.

The above reactive monomers are characteristically polymerized under conventional polymerization conditions with a hydrophilic modifier of formulae A-C or a mixture of a hydrophilic modifier of formulae A-C and a copolymerizable monomer. In those vinylic macromers containing but one vinyl group, a minor amount e.g. from about 0.01 to about 5 weight percent, based on the macromer of formula I-III, of a conventional crosslinking agent, may be employed. Suitable crosslinking agents include diolefinic monomers such as:

Allyl acrylate and methacrylate, alkylene glycol and polyalkylene glycol diacrylates and dimethacrylates, such as ethyleneglycol dimethacrylate, diethylene glycol dimethacrylate, and propylene glycol dimethacrylate; trimethylol propane triacrylate; pentaerythritol tetraacrylate, divinylbenzene; divinyl ether; divinyl sulfone; bisphenol A diacrylate or dimethacrylate; methylene bisacrylamide; diallyl phthalate; triallyl melamine and hexamethylene diacrylate and dimethacrylate. Also, such minor amounts of a crosslinking agent may be employed, if desired, in the polymerization of the di-vinyl monomer of formula I, II or III.

When the macromers of formula I-III have free hydroxy, isocyanato, carboxylic or amine groups, suitable crosslinking agents contain di- or polyfunctional co-reactive groups to form addition or condensation reactions linking 2 or more chains.

If desired, the macromer reaction mixture may contain a catalytic amount of a conventional catalyst, preferably a free radical catalyst. Of particular interest are conventional peroxide and azo catalysts, such as hydrogen peroxide, benzoyl peroxide, tert-butyl, peroctoate, benzoyl peroxide or azobis (isobutyronitrile).

The aforementioned reactions are generally straight forward additions or condensations and are typically conducted at a reaction temperature between about -10 °C and about 100 °C, depending upon the relative reactivity of the species involved, in the presence or absence of an inert diluent and in the optional presence of an addition or condensation catalyst if desired or appropriate. For reactions involving an isocyanate or acid halide, with a diol, for example, suitable optional catalysts include pyridine and triethylamine.

The siloxanes of the formula HO-Sil-H are known in the art and many are commercially readily available.

Also, the polyoxyalkylene diols of the formula HO-A-H are known or can be prepared by known methods.

Thus, the polyols of the formula HO-A-H are generally prepared by the addition reaction of t moles of an epoxide of formula X, or, assuming that A is a block copolymer of

$$CH_3$$
  
 $+(-CH_2CH-O)_{(t4)}$  and  $-(CH_2CH_2-O)_{(5)}$ 

(especially when  $\{(t_k) + (t_5)\}(t_6) = \text{about 30-70}$ ) of  $(t_k)$  time  $(t_6)$  moles of an epoxide of the formula

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$$R_7$$
 $R_8$ 
 $R_8$ 
 $R_7$ 
 $R_7$ 
 $R_7$ 
 $R_7$ 
 $R_7$ 
 $R_7$ 

with (t<sub>5</sub>) times (t<sub>6</sub>) of a different epoxide of formula X, where R<sub>7</sub>, R<sub>8</sub>, r, t, (t<sub>4</sub>), (t<sub>5</sub>) and (t<sub>6</sub>) are as defined above, optionally in the presence of a conventional alkylation catalyst, at atmospheric to elevated pressures of up to about 3000 kPa gauge, at temperatures between 0°C and about 130°C, optionally in the presence of an inert diluent. If desired, one may add to the reaction mixture, prior to the reaction of the epoxides, an aliphatic, aromatic or cycloaliphatic alcohol, acid or amine having up to 14 carbon atoms to prepare the

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The reaction between the epoxides, when mixtures of different epoxides are employed to obtain the polyol of the formula HO-A-H, can be conducted by admixing the epoxides to obtain random copolymers or terpolymers, etc., or the addition can be conducted sequentially to form block copolymers having terminal hydroxy groups. Suitable catalysts include alkaline earth oxides, alkaline earth carbonates, alkyl zinc compounds, aluminam alkoxides, hydrates of ferric chloride, bromide and acetate, and gamma radiation. The reaction may also be initiated by the presence of a glycol, such as ethylene glycol or propylene glycol or by a polyol of higher functionality such as sucrose, or by an amine, such as ethylene diamine, toluenediamine, and so forth. Generally the length of time of the reaction will depend in part on the alkylene oxide employed, but can generally be from less than one to several score hours. Thus, ethylene oxide generally is about three times as active as propylene oxide, which in turn reacts more rapidly than 1,2-butylene oxide. The preparation of polyoxetanes and polytetrahydrofurans is generally initiated via ring opening oxonium formation using trialkyloxonium salts, carboxonium salts, acylium salts and the like.

Suitable diols of the formula HO-A-H include those prepared from epoxides such as: 1,2-propylene oxide; 1.2-butylene oxide; 1,2-epoxydecane; 1.2-epoxydodecane; 1,2-epoxyoctane; 2,3-epoxynorbornane; 1,2-epoxy-3-ethoxypropane; 1,2-epoxy-3- phenoxypropane; 2,3-epoxypropyl 4-methoxyphenyl ether; tetrahydrofuran; 1,2-epoxy-3-cyclohexyloxypropane; oxetane; 1,2-epoxy-5-hexene; 1.2-epoxyethylbenzene; 1,2-epoxy-1-methoxy-2-methylpropane; perfluorohexylethoxypropylene oxide; benzyloxypropylene oxide, and the like. Also, the aforementioned epoxides may be employed as mixtures thereof. Further, certain cyclic ethers of formula X where t is 3 and the carbocyclic portion of the ring is substituted are resistant to polymerization alone, but copolymerize quite readily with more reactive cyclic ethers. Suitable comonomers include, for example, 2-methyl-tetrahydrofuran and 3-methyl-tetrahydrofuran. Also, while ethylene oxide may be employed as a co-monomer, ethylene oxide polymers, in the absence of more hydrophobic units, is characteristically too hydrophilic and absorbs too much aqueous fluid to be of use in accordance with the instant invention. However, ethylene oxide/propylene oxide copolymeric diols wherein there is 30-80 %, preferably greater than 50 %, more preferably greater than 66 % propylene oxide, on a mole basis, is sufficiently hydrophobic so as to be substantially non-swellable in aqueous media, and yet sufficiently hydrophilic so as to exhibit a contact angle with water of less than 60°; preferably less than 40°, more preferably less than 25°, more preferably less than 15°, most preferably less than 10°.

Many polymer diols of the formula HO-A-H are commercially available. Thus, suitable diol products include poloxamers having the general formula HO(CH<sub>2</sub>CH<sub>2</sub>O)<sub>B</sub> -(CH(CH<sub>3</sub>)CH<sub>2</sub>O)<sub>b</sub> -(CH<sub>2</sub>CH<sub>2</sub>O)<sub>c</sub> -H

wherein b has a value between about 16 and 100 and the sum of a and c is between about 4 and about 100. While each of the following poloxamers is beyond the current definition of the preferable A groups, they may be used to form the non-referable A groups within any one macromer. For example, if 30 % of the A groups in a macromer are poloxamers of (a) = 5, (b) = 15, (c) = 5, then the remainder of the A groups in that macromer may be any of the poloxamers set forth below. Examples of such poloxamers and their average values of a, b and c, include poloxamer 101 (a is 2, b is 16, c is 2); poloxamer 122 (a is 5, b is 21, c is 5); poloxamer 181 (a is 3, b is 30, c is 3); poloxamer 212 (a is 8, b is 35, c is 8); poloxamer 231 (a is 6, b is 39, c is 6); poloxamer 282 (a is 10, b is 47, c is 10); poloxamer 333 (a is 7, b is 54, c is 7); poloxamer 401 (a is 6, b is 67, c is 6).

Such poloxamers are available, e.g. from BASF Wyandotte under their Pluronic® brand name. Also suitable are the "reverse poloxamers", having polyethylene glycol bounded on each side by polypropylene glycol.

Polypropylene ether glycols include commercially available products having a molecular weight range between about 400 and about 4,000.

The hydrophilic modifiers can be prepared as disclosed in EP-A-331633 entitled HYDROPHILIC MODIFIER MONOMERS, invented by Frank Molock and Richard Robertson.

Typically, conventionally fully swollen polyhydroxyethyl methacrylate lenses which are spanngly crosslinked possess a  $Dk(x10^{-10})$ , (mm ml  $O_2/cm^2$  sec mmHg) value of about 5-7.

The instant polymers for use as an ophthalmic device, such as a contact lens, possess a  $Dk(x10^{-10})$  value generally greater than 7 - 10, preferably greater than about 15, more preferably greater than about 20 and still more preferably greater than about 40, yet more preferably greater than 80, even more preferably greater than 100, most preferably at least 150.

The following examples are for illustrative purposes and are not to be construed as limiting the invention. All parts are by weight unless otherwise specified.

In the examples the following test methods were used in evaluating the polymer films:

- a) Oxygen permeability (Dk) is determined from measurements on a Schema Versatae or Createch polarographic cell.
- b) Wettability or lubricity is evaluated by a feel comparison to crosslinked polyHEMA (i.e. polymerized 2-hydroxyethyl methacrylate + 1 % ethylene glycol dimethacrylate). The surface lubricity of polyHEMA is designated (B); more wettable than polyHEMA is designated (A) and a less wettable surface than polyHEMA is designated (C).
- c) Tensile strength is qualitatively evaluated by pulling on a sheet of the material by hand and then comparing the result to a material previously evaluated by a micromechanical tester to be 14 kg/cm². A measurement found to be similar to this "standard" is designated (2); a higher value is designated (1); a lower value is designated (3); while a much lower value is designated (4).

# Example 1:

This example illustrates the sequential synthesis of a polydimethyl-siloxane/polyether urethane-linked adduct.  $Z_1$  (IPDI-PEG-IEM)2, where  $Z_1$  is a polydimethyl siloxane, IPDI is isophorone diisocyanate, PEG is a polyethylene glycol and IEM is isocyanatoethyl methacrylate; and, the solution polymerization of this adduct.

# Synthesis of Z: {IPDI-PEG-IEM}2, Diadduct:

A stock solution of Z<sub>1</sub> (IPDI)<sub>2</sub> is first formed by combining 8.0054 g (0.0036 mole) Shin-Etsu X-61-504A (1000) (MW = 2212) with 1.6010 g (0.0072 mole) IPDI and 1 microliter of dibutyltin dilaurate (DBTL) and then stirring overnight at room temperature. The percentage of isocyanate reacted (generally 48-50 %) is determined by dibutylamine titration and varies with the precise value of Z<sub>1</sub> molecular weight actually used. Then, a calculated molar amount of PEG is dissolved in solvent and the corresponding amount of the Z<sub>1</sub> (IPDI)<sub>2</sub> stock solution is added to form Z<sub>1</sub> (IPDI-PEG)<sub>2</sub>. The solution is reacted at 50°C with stirring overnight. IR verifies complete reaction of the isocyanate functionality (disappearance of absorption at 2270 cm<sup>-1</sup>).

The calculated molar amount of IEM is added to a solution  $Z_1$  (IPDI-PEG)<sub>2</sub> and reacted at room temperature overnight with stirring.

# Polymerization of Diadduct; Hydrogel Formation:

The Z<sub>1</sub> (IPDI-PEG-IEM)<sub>2</sub> adduct in solvent is mixed with initiator (benzoin methyl ether, BME) poured into flat "sandwich" molds and polymerized with UV light (Sylvania, Blacklight Blue, 15 W) at room temperature for 3 hours. The polymerized films are removed from the molds, extracted in water several times, heated in boiling water for 10 minutes and then allowed to equilibrate in room temperature water (generally overnight).

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## Example 2:

This example shows the resultant water contents and oxygen permeability values of polydimethyl siloxane PEG hydrogels, i.e. crosslinked Z<sub>1</sub> (IPDI-PEG-IEM)<sub>2</sub>, solution polymerized using different molecular weights of PEG (200, 400, 600, 900, 1000, 1500, and 2000). Different solvents are used as indicated below to achieve solubility of the diadduct. Films are prepared as in example 1 except for samples 1, 2 and 3 in which PEG and Z-(IPDI)<sub>2</sub> are reacted at room temperature and samples 6 and 7 in which PEG and Z-(IPDI)<sub>2</sub> are reacted at 50° C in the presence of additional dibutyltin dilaurate catalyst.

10	Sample	Wt.% In	the Polyme	rization Mixture		<del></del>	
		Z٠	IPDI	PEG (MW)	IEM	Solvent	ВМЕ
	1	34.4	6.9	6.1 (200)	4.9	47.7 (1)	0.18
15	2	28.0	5.6	10.9-(400)	4.2	51.2 (2)	0.12
	3	27.5	5.5	15.0 (600)	3.8	48.1 (2)	0.11
	4	24.1	4.8	19.5 (900)	3.5	48.1 (1)	0.10
	5	23.0	4.6	20.7 (1000)	3.2	48.5 (1)	0.14
	6	18.8	3.8	25.5 (1500)	2.7	49.3 (3)	0.10
20	7	16.6	3.3	29.6 (2000)	2.4	48.0 (3)	0.49
	solvents:	1) = isopro	pyl acetate.	(2) = dichlorom	ethane, (3)	= acetone	
	Sample	Physical	Properties o	Hydrogel			
25		% H₂O Content	Clarity	Wettability	DK (barrier)		
	1	5.6	Clear	С	•		
	2	16.6	Clear	В	70.2	İ	I
	3	27.9	Clear	В		ļ	Ī
30	4	39.8	Clear	Ā	66.1	- 1	l
	5	46.0	Clear	A	59.5	Ī	ı
	6	57.9	SI. Hazy	A	45.2		1
	7	79.0	Opaque	A	47.5		

As shown above water content increases with higher molecular weight PEG. Oxygen permeability however, decreases with the higher molecular weight PEG.

# 40 Example 3:

This example illustrates how the clarity of polydimethyl siloxane/PEG hydrogels is affected by varying the proportion of solvent. Films are made as in example 1 except that the molecular weight of PEG is 1000 and the solvent used is isopropyl acetate (IPAC). Samples 6 and 8 are made as in example 1 with 50 % IPAC and are subsequently diluted to 60 and 75 % IPAC respectively prior to the addition of BME and UV irradiation. Samples 1 and 2 do not contain enough solvent to dissolve the reactants and, therefore, the synthesis is stopped prior to IEM addition and films are not made.

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Sample	Wt.%	in th	e Poly	meriz	ation Mix	ture	Clarity
	Zı	IPDI	PEG	IEM	Solvent	вме	
1	37.7	1	34.1	1.	20.56	-	(Not enough solvent to dissolve reactants, no films made
2	37.0	7.4	33.4	١ -	22.19	-	Same as in sample 1
3	31.8	6.4	28.5	4.4	28.91	0.54	Hazv
4	26.7	5.4	24.2	3.8	40.04		Slightly Hazy
5	23.0	4.6	20.7	3.2	48.45	0.14	
6	17.8	3.6	16.0	2.5	60.18	0.20	
7	15.2	3.0	13.7	2.2		0.12	
В	11.1	2.2	10.0	1.5		0.18	· ·

15 As shown above clarity increases with increased solvent concentration.

## Example 4:

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This example shows the effect of comonomer addition of the physical properties of Z<sub>1</sub> PEG hydrogel films. Films are made as in example 3 (sample 5) except that a comonomer (N,N-dimethyl acrylamide, DMA; N-vinyl pyrrolidone, NVP; 2-hydroxyethyl methacrylate, HEMA; or methyl methacrylate, MMA) is

Sample	Wt.%	In the Po	olymeriza	ation Mix	ture			
	Zı	IPDI	PEG	IEM	IPAC	Comonomer	BME	% H <sub>2</sub> O Content in Hydrogel
1 2 3 4 5	21.8 23.0 21.5 21.8 21.8	4.4 4.6 4.3 4.4 4.4	19.7 20.7 19.3 19.6 19.7	3.0 3.2 2.9 3.0 3.0	45.9 48.5 45.2 45.8 45.9	5.2 MMA 0 6.7 DMA 5.5 NVP 5.2 HEMA	0.10 0.14 0.13 0.15 0.13	43.7 45.5 50.0 45.7 44.4

MMA = methylmethacrylate:

DMA = N.N-dimethylacrylamide;

NVP = N-vinylpyrrolidone;

HEMA = 2-hydroxyethyl methacrylate.

As shown above, addition of comonomers has little effect on wettability or clarity. All hydrogels are clear and have an A-wettability.

# Example 5:

This example compares the physical properties of Z<sub>1</sub>/PEG hydrogel films made by combining PEG MW 1000 with lower molecular weight PEGs (200, 400 and 600) in various ratios. Except for sample 6, films are prepared as described in example 1 using isopropyl acetate (IPAC) as solvent and combining the molecular weight PEGs indicated below. Sample 6 is prepared by separately synthesis of Z<sub>1</sub>/PEG 400 and Z<sub>1</sub>/PEG 1000 according to example 1 using isopropyl acetate as solvent and then combining them prior to the addition of Initiator and UV irradiation.

EP 0 395 583 A2

	Sample	Mole Ratio	Wt.%	In the Polymeri	zation Mixture			
_		Low MW PEG:PEG 1000	Z,	IPOI	PEG (MW)	IEM	IPAC	BME
5	1	0.1	23.0	4.6	20.7 (1000)	1 20		
	2	PEG 400:PEG 1000	24.3	4.8	0.5 (200)	3.2 3.5	48.5	0.14
	1	1:7			19.2 (1000)	3.5	47.6	0.13
	3	PEG 400:PEG 1000	23.8	4.7	0.8 (400)	3.3	48.0	0.14
10	1.	1:9	1	İ	19.3 (1000)	0.0	40.0	0.14
	4	PEG 400:PEG 1000	24.2	4.9	2.2 (400)	3.4	48.8	0.14
	5	1:3			16.5 (1000)	<b>3</b>	10.0	0.14
	1 3	PEG 400:PEG 1000 1:1.5	25.5	5.1	3.7 (400)	3.7	48.2	0.16
	6	PEG 400:PEG 1000			13.8 (1000)			
15		1:1.5	25.7	5.1	3.7 (400)	3.6	49.0	0.13
	7	PEG 600:PEG 1000	24.0	1	12.9 (1000)		ļ	- 1
		1:3	24.0	4.8	3.3 (600)	3.4	48.3	0.18
	Cample		L	<u> </u>	16.2 (1000)	- 1	}	1
20	Sample	Physical Properties of	Hydrogel			1	1	
		% H₂O Content	Clarity	Wettability	Strength		-	
	1	45.4	Clear	Α	3		<del></del>	
	2	39.9	Clear	A	4	- }	- 1	
25	3 4	41.5	Clear	A	-3	- 1	- 1	ì
23	5	38.2	Clear	Α	2	1		1
1		33.4 39.8	Clear	A	2-3	- 1	1	- 1
- 1	6 7	38.5	Clear	A	3-4	j	- 1	
į		30.3	Clear	Α	4	- 1		

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The results show that incorporation of low molecular weight PEGs decreases water content but maintains clarity and wettability of the resultant films.

# Example 6:

This example demonstrates the effect of initiator level on the physical properties of Z-PEG hydrogel films with similar compositions. The films are prepared as described in example 3 (sample 5).

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Sample   Wt.% In the Polymerization Mixture   Z:   IPDI   PEG   IEM   IP.	
1 23.0 4.6 20.8 3.2 48	AC   BME
	.4 0.04
2 22.9 4.6 20.7 3.2 48	
3 23.1 4.6 20.7 3.1 48	
4 23.0 4.6 20.7 3.2 48.	
5 23.0 4.6 20.8 3.2 48.	
6 22.9 4.6 20.7 3.2 48.	6 0.16
7 22.9 4.6 20.7 3.2 48.	6 0.47
8 23.0 4.6 20.8 3.2 48.	4 0.49
9 22.9 4.6 20.7 3.2 48.	
10 23.0 4.6 20.8 3.2 48.	
Sample Physical Properties of Hydrogel	
% H <sub>2</sub> O Clarity Wettability Strength	
% H <sub>2</sub> O Clarity Wettability Strength  1 45.4 Clear A 4 2 43.4 Clear A 4	-
% H2O     Clarity     Wettability     Strength       1     45.4     Clear     A     4       2     43.4     Clear     A     4       3     46.4     Clear     A     3	
% H2O     Clarity     Wettability     Strength       1     45.4     Clear     A     4       2     43.4     Clear     A     4       3     46.4     Clear     A     3       4     45.5     Clear     A     3	
% H2O     Clarity     Wettability     Strength       1     45.4     Clear     A     4       2     43.4     Clear     A     4       3     46.4     Clear     A     3       4     45.5     Clear     A     3       5     46.3     Clear     A     3-4	
% H2O     Clarity     Wettability     Strength       1     45.4     Clear     A     4       2     43.4     Clear     A     4       3     46.4     Clear     A     3       4     45.5     Clear     A     3       5     46.3     Clear     A     3-4       6     42.7     Clear     A     3-4	
% H2O     Clarity     Wettability     Strength       1     45.4     Clear     A     4       2     43.4     Clear     A     4       3     46.4     Clear     A     3       4     45.5     Clear     A     3       5     46.3     Clear     A     3-4       6     42.7     Clear     A     3-4       7     45.7     Clear     A     2	
% H2O     Clarity     Wettability     Strength       1     45.4     Clear     A     4       2     43.4     Clear     A     4       3     46.4     Clear     A     3       4     45.5     Clear     A     3       5     46.3     Clear     A     3-4       6     42.7     Clear     A     3-4       7     45.7     Clear     A     2       8     45.9     Clear     A     2	
% H2O     Clarity     Wettability     Strength       1     45.4     Clear     A     4       2     43.4     Clear     A     4       3     46.4     Clear     A     3       4     45.5     Clear     A     3       5     46.3     Clear     A     3-4       6     42.7     Clear     A     3-4       7     45.7     Clear     A     2	

As shown above, increasing initiator level to at least 0.47 % improves the strength of the resultant hydrogels.

# Example 7:

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This example shows the effect of chain extension of the polydimethyl siloxane ( $Z_1$ ), prior to synthesis of the diadduct, on the resultant hydrogels.

3.0068 g (0.0014 mole) Shin-Etsu X-61-504A (1000) (MW 2212) are combined with 0.1512 g (0.0007 mole) IPDI and 0.5 microliters of DBTL and stirred overnight at room temperature. IR indicates complete reaction of the isocyanate functionality. This chain extended Z<sub>1</sub> is subsequently capped with IPDI and PEG as described in example 1 using PEG 1000 except for (sample 3). Sample 3 is heated at 50°C for 72 hours, (rather than overnight) during the reaction with PEG. The calculated amount of IEM is then added to form (Z<sub>1</sub>-IPDI-Z<sub>1</sub>)-(IPDI-PEG-IEM)<sub>2</sub>. The IEM is reacted at room temperature overnight with stirring except for sample 1 which is heated at 35°C.

Sample	Wt.% In	the Polymerization	n Mixture			
	Z·	IPDI	PEG 1000	IEM	Solvent	BME
.1 2 3 4	25.6 30.1 24.4 30.4	3.8 4.6 3.7 4.6	11.6 13.6 11.1 13.5	1.8 2.1 1.8 2.1	57.1 (1) 49.6 (2) 59.0 (2) 49.4 (3)	0.12 0.14 0.16 0.10
		pyl acetate, (2) =		methyl eth	yl ketone.	
Sample	Physical	Properties of Hyde	rogel			
	% H₂O Content	Clarity	Wettability	Strength	DK (barrer)	
1 2 3	40.8 36.4 43.1	Opaque Hazy Very SI. Hazy	A . A	1-2 1-2 1-2	71.0 105.2	<del></del>

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## Example 8:

This example demonstrates the fabrication of prototype polydimethylsiloxane/polyether lenses from polypropylene contact lens molds. The reaction solution is made as in example 3 (sample 5) using Shin-Etsu X-61-504A (1000) MW = 2400 and 0.78 % BME. The solution is pipetted into the lens molds and UV cured for 3 hours. After extraction in water and boiling, optically clear, strong lenses are obtained. Micromechanical tensile tests performed on the lenses yield the following properties; stress of 3.2 kg/cm², Young's modulus 15.9 kg.cm² and a 25 % elongation.

## Claims

1. A macromer comprising a segment, Seg. of the formula  $\{R \cdot a - Sil - \{(L)_h - R_1 \cdot a - Sil\}_n - \{(L)_h - R_1 \cdot a - Sil\}_n \}$  (I) wherein  $R \cdot a$  is a bond or  $-N(R_1)$ -, or if  $R \cdot a$  is not attached to a

wherein  $R_{\cdot \bullet}$  is a bond or  $-N(R_1)$ -, or if  $R_{\cdot \bullet}$  is not attached to a silicon atom, then  $R_{\cdot \bullet}$  may also be -0-;  $R_{\cdot \circ}$  is -0- or -N( $R_{\cdot}$ )-;

R. is hydrogen, C. -calkyl, or phenyl;

40 n is zero to 3:

m is zero to 3;

h is 1. or h may be zero when the atom of Sil to which it would otherwise be attached is carbon; each Sil is independently of the formula

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each L is independently of the formula  $-L_1 R_6 L_2$ -

and each A is independently of the formula

$$\begin{bmatrix}
R_7 & R_8 & R_7 \\
C & (C)_r & C & C
\end{bmatrix}$$

$$\begin{bmatrix}
R_7 & R_8 & R_7 \\
C & (C)_r & C
\end{bmatrix}$$

$$\begin{bmatrix}
R_7 & R_8 & R_7 \\
C & (C)_r & C
\end{bmatrix}$$

$$\begin{bmatrix}
R_7 & R_8 & R_7 \\
R_7 & R_8 & R_7
\end{bmatrix}$$

$$\begin{bmatrix}
R_7 & R_8 & R_7 \\
R_7 & R_8 & R_7
\end{bmatrix}$$
(V)

wherein within formula IV each R₂ and each R₅ is independently branched C₂-₂alkylene, carbonyl. C₁-₅alkylene carbonyl. C₁-₅alkylene-(oxy-C₁-₄alkylene)₀- or C₁-₄alkylene-OCH₂-(hydroxy C₁-₄alkylene)-CH₂-;

with g being an integer of from 1 to 10:

each b and each f is independently 0 or 1;

each y is independently 1 to 200;

and each R<sub>3</sub> and each R<sub>4</sub> is independently selected from C<sub>1-18</sub>alkyl and aryl of up to 12 carbon atoms; and within L:

each L1 and L2 is independently

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- CNH- in which the oxygen or nitrogen is bound to Rs. or

- C - or a bond; and

each R<sub>6</sub> is independently

(i) -(C1-25 aliphatic)- or

(ii) a group of the formula

-(C<sub>1</sub>-2<sub>0</sub> aliphatic)<sub>1</sub>-Q<sub>k</sub>-((5-7 membered cycloaliphatic or aryl of up to 25 carbon atoms)-Q<sub>(k1)</sub>-(C<sub>1</sub>-2<sub>0</sub> aliphatic)-(<sub>11</sub>)<sub>0</sub>- (IX),

each of j, k, k1 and j1 being independently zero or one, but if j is 0, then k is 0 and if j1 is 0, then k1 is 0; each D is 1 to 3;

each aliphatic group in the  $R_6$  (i) or (ii) above is uninterrupted or interrupted at any point by Q; each aliphatic, cycloaliphatic, and aryl group within the  $R_6$  (i) or (ii) above is unsubstituted or substituted by one or more substituents which substituents are halogen, hydroxy,  $C_{1-4}$ alkyl, carboxy or  $C_{1-12}$ perhaloalkyl; each Q is independently selected from

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and within formula V:

each r is independently 0-4;

each t is independently 3-200; and

each R<sub>7</sub> is independently selected from groups i) and iii) below and each R<sub>8</sub> is independently selected from groups i)-iv) below:

i) hydrogen, halogen, unsubstituted  $C_{1-16}$ alkyl, substituted  $C_{1-16}$ alkyl, unsubstituted  $C_{2-16}$ alkenyl, wherein a) the substituents on the alkyl and alkenyl groups in i) are independently selected from fluoro,  $C_{1-16}$ alkoxy,  $C_{2-16}$  alkanoyloxy,  $C_{1-16}$ alkoxy-C(0)-,  $C_{2-16}$ alkenyloxy-C(0)-,  $C_{3-16}$ alkenoyloxy, aryl of up to 10 carbon atoms, aryloxy of up to 10 carbon atoms, aroyloxy of up to 11 carbon atoms, aryl (of up to 10 carbon atoms)-oxycarbonyl,  $C_{3-8}$ cycloalkyl,  $C_{3-8}$ cycloalkoxy,  $(C_{3-8}$ cycloalkyl)-carbonyloxy,  $(C_{3-8}$ cycloalkoxy)carbonyl, oxacycloalkyl of up to 7 carbon atoms, oxacycloalkoxy of up to 7 carbon atoms, oxacycloalkyl (of up to 7 carbon atoms)-carbonyl, and oxacycloalkyl (of up to 7 carbon atoms)-carbonyloxy; and b) each alkyl and alkenyl group in a) being unsubstituted or further substituted by fluoro,  $C_{1-6}$ alkyl, or  $C_{1-6}$ alkoxy provided that said last mentioned alkoxy is not bound to a carbon already singly bound to another oxygen atom; and

ii)  $C_{1-16}$  alkoxy,  $C_{2-16}$  alkanoyloxy,  $C_{1-16}$  alkoxy-C(0)-,  $C_{2-16}$  alkenyloxy-C(0)- and  $C_{3-16}$  alkenoyloxy, each of which may be further substituted by fluoro, aryl of up to 10 carbon atoms, or  $C_{1-6}$  alkoxy provided that the last mentioned alkoxy is not bound to a carbon atom which is singly bound to another oxygen;

iii) aryl of up to 10 carbon atoms, C<sub>3-8</sub>cycloalkyl, and oxacycloalkyl of up to 7 carbon atoms, each of which is unsubstituted or substituted by a substituent selected from those indicated in i)a) and b) above;

iv) aryloxy of up to 10 carbon atoms, cycloalkoxy of up to 8 carbons, cycloalkyl (of up to 8 carbons)-C(O)O-, cycloalkoxy (of up to 8 carbons)-carbonyl, aroyloxy of up to 11 carbons, oxacycloalkoxy of up to 7 carbons, oxacycloalkenyloxy of up to 7 carbons, oxacycloalkenyloxy of up to 7 carbon atoms)-carbonyloxy, aryloxy of up to 11 carbon atoms, and aroyloxy of up to 11 carbon atoms, and aroyloxy of up to 11 carbon atoms, each of which is unsubstituted or substituted by fluoro,  $C_{1-\epsilon}$  alkyl, or  $C_{1-\epsilon}$  alkoxy, provided said last mentioned alkoxy is not bound to a carbon atom which is singly bound to another oxygen; and 2 adjacent groups  $R_7$  and  $R_8$ , together with the atoms to which they are attached may define a 5-8 membered cycloalkyl, cycloalkenyl, or oxacycloalkyl ring or a 6-14 membered bicyclo-ring.

2. The macromer of claim 1 comprising a segment, Grp. of the formula

 $\{R_{15}-A-\{L-R_{15}-A-\}_m (L)_h\}_a$   $\{Seg-(L)_h\}_p-Seg-((L)_h-R_{14}-Sil-[(L)_h-R_{14}-Sil-]_n\}_e$  (II) wherein  $R_{14}$ ,  $R_{15}$ , L, Seg, Sil, h and A are as defined in claim 1;

m is an integer of zero to 10; n is an integer of zero to 3;

p is an integer of zero to 7; and

a and e are each zero or 1, but a plus e is 1.

3. The macromer of claim 2 of the formula

Cap-Grp-Cap III

wherein Grp is as defined in claim 2:

each Cap is independently selected from formula VI or VII below provided at least one Cap per macromer is of formula VI; wherein formula VI is

 $R_{16}(L_4)_x(R_6)_x(A_1)_x(L_1)_x(J(R_9)_M(L_5)_G(R_6)_x(L_6)_T]_{\overline{z}}$  (VI)

and formula VII is

 $R_{17}(L_4)_z(R_6)_w(A_1)_x(L_3)_x(J(R_9)_w(L_5)_G(R_6)_x(L_6)_7$  (VII)

in which each Rs is as defined in claim 1;

each w, x and z is independently zero or 1, but if w is zero, then one of x and z must be zero; each G, M, T, V, Y and Z is independently zero or 1, but if Y is 0, then G must be zero and M must be one; each J is -O- or -NH-;

each A<sub>1</sub> and R<sub>2</sub> is independently of the formula

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$$\begin{bmatrix}
R_7 & R_8 & R_7 \\
C - (C)_r - C - C - C \\
R_7 & R_8 & R_7
\end{bmatrix}_{1-7}$$
;

each La is independently

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with the right hand portion attached to Grp provided that the Grp atom to which L<sub>3</sub> is attached is not a carbonyl carbon, an amido nitrogen or a carboxy oxygen and L<sub>3</sub> is -O- or -NR<sub>1</sub>- when such Grp atom is a each L is independently

when w is one, and selected from the same group as L3 when w is zero; each Ls is -C(0)NH- with the carbonyl attached Rs to J: each  $L_{\epsilon}$  is -C(0)NH- with the nitrogen attached to  $R_{\epsilon}$  or  $R_{9}$ ; each Ris is independently of the formula

(VIII)

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in which  $R_{12}$  is hydrogen or  $C_{1-7}$ alkyl and each of  $R_{13}$  and  $R_{11}$  is independently hydrogen,  $C_{1-7}$ alkyl, or -COOR13 with R13 being hydrogen or C1-4alkyl; and Rin is selected from

- a) H, NH2, OH, -NCO, -NCS, -C(O)-halogen and COOR12 when the non-R17 atom to which it is attached is a non-carbonyl saturated carbon or the non-R17 atom to which it is attached is a carbon which is part of a ring;
- b) H, -C(O)-halogen, -C(O)-R<sub>12</sub>, -C(O)-OR<sub>12</sub> and R<sub>12</sub> when the non-R<sub>17</sub> atom to which it is attached is an oxygen other than a carboxy oxygen:
  - c) H and R-2, when the non-R17 atom to which it is attached is a carboxy oxygen;
- d) -C(0)-R-2, -C(0)-OR12, -C(0)N(R1)R12 and R-2 when the non-R17 atom to which it is attached is an amino nitrogen, or R12 when the non-R17 atom to which it is attached is an amido nitrogen;
- e) OH, R:2, CH2C(O)R:2, CH2C(O)OR:2, NH2, and -NHR:2 when the non-R:7 atom to which it is attached is a carboxy carbon: and
  - f) H, OH, hal, -OR12 when the non-R17 atom to which it is attached is silicon.
- 4. A macromer according to any of claims 1 to 3 wherein at least 30 % of the A blocks within any one macromer of formulae I-III are homopolymeric for not more than 15 repeating units.
- 5. A macromer according to claim 1 wherein Seg is represented by Segi of formula {R·+-Sil-(L)h-R+5-A} (la)
- wherein R14, R15, Sil, L, A and h are as defined in claim 1.
- 6. A macromer according to claim 2 wherein Grp is represented by Grp1 of formula  $(R_{15}-A-(L)_h)_a-Seg:-((L)_h-R_{14}-Sil)_e$ (IIa) wherein R. a., R. S. A. L. Sil, a. e and h are as defined in claim 2 and Seg. is as defined in claim 5.
- 7. A macromer according to claim 3 wherein Grp is represented by Grp, as defined in claim 6 and Cap is represented by Cap<sub>1</sub> of formulae  $R_{16}(L_4)_z(R_6)_w(A_1)_v(L_3)_x$ (Vla) and

 $R \cdot r(L_4)_z(R_6)_w(A_1)_v(L_3)_{\overline{x}}$ (VIIa)

wherein all the variables are as defined in claim 3.

- 8. A macromer according to any of claims 5-7 wherein at least 30 % of the A blocks within any one macromer of formulae la, lla or Illa are homopolymeric for not more than 15 repeating units.
  - 9. A macromer according to claim 3 wherein

CH<sub>3</sub> CH<sub>3</sub> CH<sub>3</sub> CH<sub>3</sub>
-CHCH<sub>2</sub>CH<sub>2</sub>—(Si-O)<sub>y</sub>—Si-CH<sub>2</sub>CH<sub>2</sub>CH-O-;
CH<sub>3</sub> CH<sub>3</sub> Sil is

a is one;

e is zero;

Ris and Ris are each -O-;

L is

10 h is one:

A is (CH2CH2O)1; and

p is zero:

m is zero:

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10. A macromer according to claim 4 wherein all the variables have the meaning as in claim 9.

11. A macromer of the formula

Cap-Sil-L-O-A-L-Sil-Cap

wherein Sil is 
$$-CH_2CH_2 - (Si-O)_y - Si-CH_2CH_2 - (CH_3)_y - CH_3$$

wherein y is 1 to 200, (especially y of about 4, 7, and about 16);

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with the phenyl ring thereof bound to Sil: m is zero to 3;

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$$CH_3$$
  
A is - $(CH_2CH_2O)_{t^-}$ , - $(CH_2CH_2O)_{t^-}$ 

wherein t is 3 to 200, or a poloxamer or reverse poloxamer having t polyoxyalkylene units per A group (especially t of about 6-8, about 45-55, and about 90-95); and

- 12. A macromer according to claim 11 wherein all the variables have the meaning as defined in claim 11 and wherein at least 30 % of the A blocks are homopolymeric for not more than 15 repeating units.
- 13. A polymer fabricated from one or more macromers of any of claims 1 to 3 having in its fully swollen state a water content of at least 10 %.
- 14. A polymer fabricated from one or more macromers of claim 4 having in its fully swollen state a water content of at least 10 %.
  - 15. A polymer fabricated from one or more macromers of any of claims 5 to 7 having in its fully swollen state a water content of at least 10 %.
- 16. A polymer fabricated from one or more macromers of claim 8 having in its fully swollen state a water content of at least 10 %.
  - 17. A polymer fabricated from one or more macromers of claim 9 or 10 having in its fully swollen state a water content of at least 10 %.
  - 18. A polymer fabricated from one or more macromers of claim 11 or 12 having in its fully swollen state a water content of at least 10 %.
- 19. A polymer being a copolymer of at least 10 % of one or more macromers of any of claims 1 to 3 and at least one copolymerizable monomer which is not a macromer of any of claim 1 to 3, having in its fully swollen state a water content of at least 10 %.
- 20. A polymer being a copolymer of one or more macromers of claim 4 and at least one copolymerizable monomer which is not a macromer of claim 4, having in its fully swollen state a water content of at least 10 %.
  - 21. A polymer being a copolymer of at least 10 % of one or more macromers of any of claims 5 to 7 and at least one copolymerizable monomer which is not a macromer of any of claim 5 to 7, having in its fully swollen state a water content of at least 10 %.
- 22. A polymer being a copolymer of one or more macromers of claim 8 and at least one copolymerizable monomer which is not a macromer of claim 8, having in its fully swollen state a water content of at least 10 %.
  - 23. A polymer being a copolymer of one or more macromers of claim 9 or 10 and at least one copolymenizable monomer which is not a macromer of claim 9 or 10, having in its fully swollen state a water content of at least 10 %.
- 24. A polymer being a copolymer of one or more macromers of claim 11 or 12 and at least one copolymerizable monomer which is not a macromer of claim 11 or 12, having in its fully swollen state a water content of at least 10 %.
  - 25. An ophthalmic device comprising an optically clear polymer of claim 13 or 14.
  - 26. An ophthalmic device comprising an optically clear polymer of claim 15 or 16.
  - 27. An ophthalmic device comprising an optically clear polymer of claim 17.
    - 28. An ophthalmic device comprising an optically clear polymer of claim 18.
    - 29. An ophthalmic device comprising an optically clear polymer of claim 19 or 20.
    - 30. An ophthalmic device comprising an optically clear polymer of claim 21 or 22.
    - 31. An ophthalmic device comprising an optically clear polymer of claim 23.
    - 32. An ophthalmic device comprising an optically clear polymer of claim 24.
    - 33. The ophthalmic device of any of claims 25, 26, 29 or 30 which is a contact lens.
    - 34. The ophthalmic device of any of claims 27 or 31 which is a contact lens.
    - 35. The ophthalmic device of any of claims 28 or 32 which is a contact lens.
- 36. A method of correcting visual defects comprising applying to a patient in need thereof a visual defect correcting opthalmic device of claim 25.
  - 37. A method of changing eye color comprising applying to a patient a colored contact lens of claim 33.

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